ITED STATES PATENT AND TRADEMARK OFFICE

Group Art Unit:

Examiner: I. MARX

In re Application of: Venkataraman BRINGI et al.

09/083,198

Filed: May 22, 1998

For:

Application Number:

ENHANCED PRODUCTION OF TAXOL AND TAXANES BY CELL CULTURES

OF TAXUS SPECIES

SUPPLEMENT TO SUBMISSION OF MARCH 19, 2002

Commissioner for Patents U.S. Patent and Trademark Office Washington, D.C. 20231

Sir:

Applicants hereby submit the attached passages from "The Merck Index, 12th edition, pages 365 and 986. These pages were inadvertently omitted from the Submission filed on Mach 19, 2002.

Applicants believe that the application is now in condition for allowance, and a Notice to that effect is earnestly solicited.

No fee is believed to be required for this submission. However, in the event any fee is deemed necessary for consideration of all materials submitted in response to the most recent Office Action, the Commissioner is authorized to charge the undersigned's Deposit Account No. 50-1640.

Respectfully submitted,

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March 25, 2002

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LHP:cdh

propyl)phenothiazine; 3-chloro-10-(3-diethylaminopropyl)phenothiazine; RP-4909. C₁₉H₂₂ClN₂S; mol wt 346.92. C 65.78%. H 6.68%. Cl 10.22%. N 8.07%, S 9.24%. Prepn: Buisson et al. U.S. pat. 2,769,002 (1956 to Rhône-Poulenc).

Hydrochloride, C₁₉H₁₃CIN₂S.HCl, Neuriplege. Crystals, mp 178°. (Free base bp₁ 225-240°). Sensitive to light. Soly in water about 1.0 g/60 ml, ethanol about 1.0 g/300 ml, ethoroform 1.0 g/5 ml. Practically insol in acetone, ether, benzene. pH of 1% aq soln 4.8.

THERAP CAT: Muscle relaxant (skeletal); antipsychotic.

2237. Chlorproguanil. N-(3,4-Dichlorophenyl)-N'-(1-methylethyl)imidodicarbonimidic diamide; 1-(3,4-dichlorophenyl)-5-isopropylbiguanide; N'-3,4-dichlorophenyl-N'-isopropylbiguanide; N'-3,4-dichlorophenyl-N'-isopropyldiguanide; N'-3,4-dichlorophenyl-N'-isopropyldiguanide; N-5943. C₁₁H₁₅Cl₂N₃; mol wt 288.18. C 45.85%, H. 5.25%, Cl 24.60%, N 24.30%. Method of prepn: Crowther et al., J. Chem. Soc. 1951, 1780; Curd et al., U.S., pat. 2,544,827 (1951); Crowther et al., Brit. pat. 667,116 (1952) (both to ICI).

Hydrochloride, $C_{11}H_{15}Cl_2N_5$.HCl, Lapudrine. Crystals, mp 246-247. Soly in water: 1 g/100 ml. Solns may be boiled without dec.

THERAP CAT: Antimalarial.

2238. Chlorpromazine. 2-Chloro-N,N-dimethyl-10H-phenothiazine: 10-propanamine; 2-chloro-10-(3-dimethylaminopropyl)phenothiazine; 3-chloro-10-(3-dimethylaminopropyl)phenothiazine; N-(3-dimethylaminopropyl)-3-chloro-phenothiazine; 2601-A; HL-5746; RP-4560; SKF-2601-A; Chlorderazin; Chlorpromados; Contomin; Esmind; Fenactil; Novomazina; Promactil; Prozil; Plegomazin; Sanopron; Aminazine; Ampliactil; Amplictil; Promazil; Proma; Elmarin; Wintermin. C₁₇H₁₉ClN₂S; mol wt 318.87. C 64.03%, H 6.01%, Cl 11.12%, N 8.79%, S 10.06%. Prepn: Charpentier et al., Compt. Rend. 235, 59 (1952); Charpentier, U.S. pat. 2,645,640 (1953 to Rhône-Poulenc). Effects of neuroleptics on dopamine receptors: N.-E. Anden et al., Eur. J. Pharmacol. 11, 303 (1970). Toxicity study: E. I. Goldenthal, Toxicol. Appl. Pharmacol. 18, 185 (1971). Review of analytical methods for determin in pharmaceutical prepns: L. F. S. Chagonda, J. S. Millership, J. Pharm. Biomed. Anal. 7, 271-278 (1989). Brief historical review: G. Curzon, Trends Pharmacol. Sci. 11, 61-63 (1990).

Oily liq. Amine odor. Alkaline reaction. bp_{0.8} 200-205°. Maleate. Clordelazin.

Hydrochloride, C₁₇H₁₉ClN₂S.HCl, Hebanil, Hibanil, Hibernal, Klorpromex, Largactil, Largaktyl, Megaphen, Promacid, Chloractil, Chlorazin, Sonazine, Marazine, Propaphenin, Taroctyl, Thorazine, Torazina. Crystals, dec 179-180' (capillary); 194-196' (microblock). uv curve: Neuhoff, Auterhoff, Arch. Pharm. 288, 400 (1955). pH of 5% aq soln

4.0-5.5. One gram dissolves in 2.5 ml water. Sol in methanol, ethanol, chloroform. Practically insol in ether, benzene. Slightly acid to litmus. LD $_{50}$ orally in rats: 225 mg/kg (Goldenthal).

THERAP CAT: Antiemetic; antipsychotic.
THERAP CAT (VET): Antiemetic; tranquilizer.

2239. Chlorpropamide, 4-Chloro-N-{(propylamino)-carbony||benzenesu||fonamide; I-{p-chlorophenylsu|fony|)-3-propylurea; 1-{p-chlorobenzenesu||fony|)-3-propylurea; N-{p-chlorobenzenesu||fony|)urea; P-607; Adiaben; Asucrol; Catanil; Chloronase; Diabechlor; Diabenal; Diabetoral; Diabinese; Melitase; Millinese; Oradian; Stabinol. C₁₀H₁₁ClN₂O₃S; mol wt 276.74. C 43.40%, H 4.73%, Cl 12.81%, N 10.12%, O 17.34%, S 11.59%. Prepn: Marshall, Sigal, J. Org. Chem. 23, 927 (1958); Brit. pat. 853,555; W. M. McLamore, U.S. pat. 3,349,124 (1960, 1967 both to Pfizer); Bauer et al., J. Org. Chem. 31, 3440 (1960). Pharmacology and metabolism: Khurana et al., Indian J. Med. 55, 1084 (1967); Brotherton et al., Clin. Pharmacol. Ther. 10, 505 (1969); Madsen et al., Eur. J. Pharmacol. 31, 374 (1971). Toxicity study: E. I. Goldenthal, Toxicol. Appl. Pharmacol. 18, 185 (1971).

Crystals from dil ethanol, mp 127-129°. uv max (0.01N HCl): 232.5 nm (e 16500). Soly in water at pH 6: 2.2 mg/ml. Practically insol at pH 7.3. Sol in alc; moderately sol in chloroform; sparingly sol in ether, benzene. LD₅₀ i.p. in rats: 580 mg/kg (Goldenthal).

THERAP CAT: Antidiabetic.

2240. Chlorpropham. (3-Chlorophenyl)carbamic acid 1-methylethyl ester; m-chlorocarbanilic acid isopropyl ester; isopropyl-m-chlorocarbanilate; isopropyl N-(3-chlorophenyl)carbamate; chloro-IPC; chloropropham; CIPC; Chlor-IFC; Furloe; Sprout-Nip. C₁₀H, CINO₂; mol wt 213.66. C 56.21%. H 5.66%. CI 16.59%. N 6.56%, O 14.98%. Prepn: E. D. Witman, U.S. pat. 2,695,225; Strain, U.S. pat. 2,734,911 (1954, 1956 both to Columbia-Southern Chem.); Brockway, U.S. pat. 2,806,051 (1957 to B. F. Goodrich). Toxicology: E. M. Boyd, E. Carsky, Arch. Environ. Health 19,621 (1969).

Solid, mp 40.7-41.1°. bp₂ 149°. n_D^{20} 1.5388. Commercial product is a liquid. Slightly sol in water; miscible with most oils and organic solvents. LD₅₀ orally in rats: 1.2 g/kg (Boyd, Carsky).

USE: Herbicide; plant growth regulator.

2241. Chlorprothixene. 3-(2-Chloro-9H-thioxanthen-9-ylidene)-N,N-dimethyl-1-propanamine; 2-chloro-N,N-dimethyl-1-propanamine; 2-chloro-N,N-dimethyl-3-thioxanthene-Δ**-propylamine; 2-chloro-9-(3'-dimethyl-aminopropylidene)thioxanthene; α-2-chloro-10-(3-dimethyl-aminopropylidene)thioxanthene; N-714; Taractan; Truxal; Truxaletten; Tarasan. C_HH₁₆CNS; mol wt 315.87. C 68.45%, H 5.74%, Cl 11.22%, N 4.43%, S 10.15%. Prepn: Brit. pat. 829,763 and Sprague, Engelhardt, U.S. pat. 2,951,082 (both 1960 to Merck & Co.); Brit. pat. 834,143 (1960 to Am. Cyanamid). Comprehensive description: B. C. Rudy, B. Z. Senkowski, in Analytical Profiles of Drug Substances vol. 2, K. Florey, Ed. (Academic Press, New York, 1973) pp 63-84.

(CH,O),C,H,COOH. The acid itself is unstable, rapidly changing to lactone.

5826. Mecoprop. (±)-2-(4-Chloro-2-methylphenoxy)propanoic acid; (±)-2-[(4-chloro-o-tolyl)oxy]propionic acid; mechlorprop; MCPP; CMPP; RD-4593; Astix CMPP; Iso-Cornox; Compitox; Compitox Plus; Proponex-Plus. C₁H₁₁: ClO₃; mol wt 214.65. C 55.96%, H 5.17%, Cl 16.52%, O 22.36%. Prepn: M. E. Synerholm, P. W. Zimmerman, Contrib. Boyce Thompson Inst. 14, 91 (1945). Studies on Plant growth regulation: C. H. Fawcett et al., Ann. Appl. Biol. 40, 231 (1953); and comparison of enantiomers: M. Matell, Kungl. Lantbruks-Hogsk. Ann. 20, 207 (1953); B. Aberg, ibid. 241. GLC determn: H. G. Higson, D. Butler, Analyst 85, 657 (1960). Crystal structure: G. Smith et al., Acta Crystallogr. B36, 992 (1980). Herbicidal activity: G. B. Lush, Proc. 3rd Brit. Weed Contr. Conf. 625 (1956); E. L. Leafe, ibid. 633; B. Wallgren, Weeds Weed Contr. 24th Swedish Weed Conf. 30 (1983); of (+)-enantiomer: J. Toll, Weeds Weed Contr. 28th Swedish Weed Conf. 100 (1987). Degradation in soils: L. Lindholm et al., Acta Agr. Scand. 32, 429 (1982); A. E. Smith, Bull. Environ. Contam. Toxicol. 34, 656 (1985). Toxicological studies: M. R. Gurd et al., Food Cosmet. Toxicol. 3, 883 (1965); H. G. Verschuuren et al., Toxicology 3, 349 (1975); R. Roll, G. Matthiaschk, Arzneimittel-Forsch. 33, 1479 (1983). EC-GLC determn in tissues and biological fluids: J. De Beer et al., Vet. Hum. Toxicol. 21, Suppl., 172 (1979). HPLC resolution of enantiomers: B. Blessington et al., J. Chromatog. 396, 177

Solid, mp 93-94°. LDsn in rats (mg/kg): 1210 orally, 402 i.p. (Verschuuren).

(+)-Form, Mecoprop-P, Duplosan KV. Solid, mp 95-96'. $[a]_{D}^{25} + 19^{\circ}$ (alcohol).

[a]j + 19 (alcono).

Sodium salt, C₁₀H₁₁ClNaO₃, LD₅₀ i.p. in rats, mice: 500, 600 mg/kg; orally in mice: 650 mg/kg (Gurd).

Diethylamine salt, C₁₁H₁₂ClNO₃, Mecopar. LD₅₀ in rats, mice (mg/kg): 1060 ±120, 600 ±35 orally; 350, 400 i.p.

Potassium salt, C10H11C1KO3, Mecomec, Hedonal MCPP. USE: Herbicide.

5827. Mecrylate. 2-Cyano-2-propenoic acid methyl ester; 2-cyanoacrylic acid methyl ester; methyl 2-cyanoacrylate; methyl α-cyanoacrylate; AD/here; Coapt. C₃H₄NO₃; mol wt 111.10. C 54.05%, H 4.54%, N 12.61%, O 28.80%. CH₂=C(C≡N)COOCH₃. Prepn: McKeever, U.S. pat. 2,-912,454 (1959 to Rohm & Haas); McKeever, Raterink, U.S. pat. 2,926,188 (1960 to Rohm & Haas).

Liquid, bp_{1.8} 47-49°. n²₁₀ 1.443. USE: Manuf of polymers and adhesives, see U.S. pats. 2,776,232 and 2,794,788 (1957 to Eastman Kodak). Surgical aid (tissue adhesive).

Mecysteine Hydrochloride. L-Cysteine methyl ester hydrochloride; methyl cysteine hydrochloride; methyl β-mercaptoalanine hydrochloride; methyl α-amino-β-mercaptopropionate hydrochloride; LJ-48; Acdrile; Visclair. C₄H₁₀ClNO₂S; mol wt 171.65. C 27.99%, H 5.87%, Cl 20.65%, N 8.16%, O 18.64%, S 18.68%. HSCH₂CH(NH₂)-COOCH, HCl. Prepn: Bergmann, Michalis, Ber. 63, 987 (1930); Zervas, Theodoropoulos, J. Am. Chem. Soc. 78, 1359 (1956).

Crystals from methanol, mp 140-141°. $[\alpha]_D^{20} = 2.9^\circ$ (methanol).

THERAP CAT: Mucolytic.

5829. Medazepam. 7-Chloro-2,3-dihydro-1-methyl-5phenyl-1H-1,4-benzodiazepine; Ansilan; Diepin; Medazepol; Megasedan; Narsis; Nobrium; Psiquium; Resmit; Rudotel; Tranquilax. C₁₆H₁₅ClN₃; mol wt 270.76. C 70.98%, H 5.58%, Cl 13.09%, N 10.35%. Prepn: L. H. Sternbach et al. J. Org. Chem. 28, 2456 (1963); G. A. Archer et al., Belg. pat. 620,773, C.A. 59, 10095b (1963); E. Reeder, L. H. Sternbach, U.S. pat. 3,243,427 (1963, 1966 both to Hoffmann-La Roche); S. Inaba et al., Chem. Pharm. Bull. 20, 1628 (1972); M. Mihalic et al., J. Heterocycl. Chem. 14, 941 (1977). Pharmacology: L. O. Randall et al., Arch. Int. Pharmacodyn. Ther. 185, 135 (1970). Crystal structure: G. Gilli et al., Acta Crystallogr. B34, 3793 (1978).

Colorless prismatic crystals from ether + 95-97°. LD₅₀ in mice (mg/kg): 360 i.p., 1070 orally (Randall)

Hydrochloride, C16H15CIN1.HCl, orange-red crystalline

powder. Freely sol in water, alcohol.

Note: This is a controlled substance (depressant) listed in the U.S. Code of Federal Regulations, Title 21 Part 1308.14 (1995).

THERAP CAT: Anxiolytic.

5830. Medetomidine. 4-[1-(2,3-Dimethylphenyl)ethyl]. 1H-imidazole; (±)-4-(\alpha,2,3-trimethylbenzyl)imidazole; 4-[(\alpha-1H-imidazole; (±)-4-(a,2,3-trimethylbenzyl)imidazole; 4-((a-methyl)-2,3-dimethylbenzyl]imidazole. C₁₁H₁₆N₂; mol wt 200.28. C 77.96%, H 8.05%, N 13.99%. a₂-Adrenergic agonist. Prepn: A. J. Karjalainen et al., Brit. pat. Appl. 2,101,114; A. J. Karjalainen, K. O. A. Kurkela, U.S. pat. 4,544,664 (1983, 1985 both to Farmos). Receptor binding study: R. Virtanen et al., Eur. J. Pharmacol. 150, 9 (1988). Sedative and cardiovascular effects in humans: M. Scheinin et al., Brit. J. Clin. Pharmacol. 24, 443 (1987). Veterinary evaluation in cats: D. Stenberg et al., J. Vet. Pharmacol. Ther. 10, 319 (1987).

Hydrochloride, C₁₃H₁₆CN₂.HCl, MPV-785, Domitor. d-Form, dexmedetomidine, (S)-medetomidine, MPV-1440. Pharmacokinetics: K. T. Kivisto et al., Eur. J. Clin. Pharmacol. 46, 345 (1994). Clinical evaluation as surgical premedicant: M. Virkkila et al., Anaesthesia 49, 853 (1994). THERAP CAT (VET): Sedative; analgesic.

5831. Medibazine. 1-(1,3-Benzodioxol-5-ylmethyl)-4-(diphenylmethyl)piperazine; 1-(diphenylmethyl)-4-piperanyl-piperazine; 1-benzhydryl-4-piperonylpiperazine. C₁₅H₂₆N₃. O₂; mol wt 386.49. C 77.69%, H 6.78%, N 7.25%, O 8.28%. Prepn: Belg. pat. 616,371; Regnier et al., U.S. pat. 3,119,-826 (1962, 1964, both to Science Union). Pharmacology: Laubic et al., Arch. Int. Pharmacodyn. Ther. 151, 313 (1964); Laubie, Schmitt, ibid. 155, 1 (1965).

Dihydrochloride, C25H26N2O2.2HCl, Vialibran. Solid. mp 288°.

THERAP CAT: Vasodilator (coronary); bronchodilator.

5832. Medicagol. 3-Hydroxy-6H-[1,3]dioxolo[5,6]benzofuro[3,2-c][1]benzopyran-6-one; 7-hydroxy-11,12-(methylenedioxy)coumestan; 7-hydroxy-5',6'-methylenedioxybenzo-